

Section II, or may be derived from shorter sequence regions within cloned cDNA fragments. Shorter fragments can be prepared by enzymatic digestion of full-length fragments under conditions which yield
5 desired-sized fragments, as will be described in Section IV. Alternatively, the fragments can be produced by oligonucleotide synthetic methods, using sequences derived from the cDNA fragments. Methods or commercial services for producing selected-sequence
10 oligonucleotide fragments are available. Fragments are usually at least 12 nucleotides in length, preferably at least 14, 20, 30 or 50 nucleotides, when used as probes. Probes can be full length or less than 500, preferably less than 300 or 200, nucleotides
15 in length.

To confirm that a given ET-NANB fragment is in fact derived from the ET-NANB viral agent, the fragment can be shown to hybridize selectively with cDNA from infected sources. By way of illustration, to
20 confirm that the 1.33 kb fragment in the pTZKF1(ET1.1) plasmid is ET-NANB in origin, the fragment was excised from the pTZKF1(ET1.1) plasmid, purified, and radiolabeled by random labeling. The radiolabeled fragment was hybridized with fractionated cDNAs from
25 infected and non-infected sources to confirm that the probe reacts only with infected-source cDNAs. This method is illustrated in Example 4, where the above radiolabeled 1.33 kb fragment from pTZKF1(ET1.1) plasmid was examined for binding to cDNAs prepared
30 from infected and non-infected sources. The infected sources are (1) bile from a cynomolgus macaque infected with a strain of virus derived from stool samples from human patients from Burma with known ET-NANB infections and (2) a viral agent derived from the
35 stool sample of a human ET-NANB patient from Mexico. The cDNAs in each fragment mixture were first amplified by a linker/primer amplification method described in Example 4. Fragment separation was on

Thirty ml of a 10% stool suspension obtained from an individual from Mexico diagnosed as infected with ET-NANB as a result of an ET-NANB outbreak, and a similar volume of stool from a healthy, non-infected
5 individual, were layered over a 30% sucrose density gradient cushion, and centrifuged at 25,000 x g for 6 hr in an SW27 rotor, at 15°C. The pelleted material from the infected-source stool contained 27-34 nm VLP particles characteristic of ET-NANB infection in the
10 infected-stool sample. RNA was isolated from the sucrose-gradient pellets in both the infected and non-infected samples, and the isolated RNA was used to produce cDNA fragments as described in Example 1.

The cDNA fragment mixtures from infected and
15 non-infected bile source, and from infected and non-infected human-stool source were each amplified by a novel linker/primer replication method described in co-owned patent application serial number 07/208,512 for "DNA Amplification and Subtraction Technique,"
20 filed June 17, 1988. Briefly, the fragments in each sample were blunt-ended with DNA Pol I then extracted with phenol/chloroform and precipitated with ethanol. The blunt-ended material was ligated with linkers having the following sequence (top or 5' sequence has
25 SEQ ID NO:21; bottom or 3' sequence has SEQ ID NO:22):

5'-GGAATTCTGGCGGCCGCTCG-3'
3'-TTCCTTAAGCGCCGGCGAGC-5'

The duplex fragments were digested with
30 NruI to remove linker dimers, mixed with a primer having the sequence 5'-GGAATTCTGGCGGCCGCTCG-3', and then heat denatured and cooled to room temperature to form single-strand DNA/primer complexes. The complexes were replicated to form duplex fragments by addition of
35 *Thermus aquaticus* (Taq) polymerase and all four deoxynucleotides. The replication procedures, involving successive strand denaturation, formation of